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(54) Title: ISOLATED, TYROSINASE DERIVED PEPTIDES AND USES THEREOF

# (57) Abstract

The invention relates to the identification of complexes of human leukocyte antigen molecules and tyrosinase derived peptides on the surfaces of abnormal cells. The therapeutic and diagnostic ramifications of this observation are the subject of the invention.

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# ISOLATED, TYROSINASE DERIVED PEPTIDES AND USES THEREOF

### RELATED APPLICATION

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This application is a continuation-in-part of Serial No. 08/203,054 filed on February 28, 1994, which is a continuation-in-part of copending application Serial No. 08/081,673, filed June 23, 1993, which is a continuation in part of copending U.S. Patent Application Serial Number 054,714, filed April 28, 1993 which is a continuation-in-part of copending U.S. patent application Serial Number 994,928, filed December 22, 1992.

# FIELD OF THE INVENTION

This invention relates to isolated peptides, derived from tyrosinase which are presented by HLA-A2 and HLA-B44 molecules and uses thereof. In addition, it relates to the ability to identify those individuals diagnosed with conditions characterized by cellular abnormalities whose abnormal cells present complexes of these peptides and HLA-A2 and HAL-B44, the presented peptides, and the ramifications thereof.

# BACKGROUND AND PRIOR ART

The process by which the mammalian immune system recognizes and reacts to foreign or alien materials is a complex one. An important facet of the system is the T cell response. This response requires that T cells recognize and interact with complexes of cell surface molecules, referred to human leukocyte antigens ("HLA"), major histocompatibility complexes ("MHCs"), and peptides. The peptides are derived from larger molecules which are processed by the cells which also present the HLA/MHC molecule. this regard Male et al., Advanced Immunology (J.P. Lipincott Company, 1987), especially chapters 6-10. The interaction of T cell and complexes of HLA/peptide is restricted, requiring a T cell specific for a particular combination of an HLA molecule and a peptide. If a specific T cell is not present, there is no T cell response even if its partner complex is Similarly, there is no response if the specific complex is absent, but the T cell is present. This mechanism

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is involved in the immune system's response to foreign materials, in autoimmune pathologies, and in responses to cellular abnormalities. Recently, much work has focused on the mechanisms by which proteins are processed into the HLA binding peptides. See, in this regard, Barinaga, Science 257: 880 (1992); Fremont et al., Science 257: 919 (1992); Matsumura et al., Science 257: 927 (1992); Latron et al., Science 257: 964 (1992).

The mechanism by which T cells recognize cellular abnormalities has also been implicated in cancer. For example, in PCT application PCT/US92/04354, filed May 22, 1992, published on November 26, 1992, and incorporated by reference, a family of genes is disclosed, which are processed into peptides which, in turn, are expressed on cell surfaces, which can lead to lysis of the tumor cells by specific CTLs. The genes are said to code for "tumor rejection antigen precursors" or "TRAP" molecules, and the peptides derived therefrom are referred to as "tumor rejection antigens" or "TRAS". See Traversari et al., Immunogenetics 35: 145 (1992); van der Bruggen et al., Science 254: 1643 (1991), for further information on this family of genes.

In U.S. patent application Serial Number 938,334, the disclosure of which is incorporated by reference, nonapeptides are taught which bind to the HLA-A1 molecule. The reference teaches that given the known specificity of particular peptides for particular HLA molecules, one should expect a particular peptide to bind one HLA molecule, but not others. This is important, because different individuals possess different HLA phenotypes. As a result, while identification of a particular peptide as being a partner for a specific HLA molecule has diagnostic and therapeutic ramifications, these are only relevant for individuals with that particular HLA There is a need for further work in the area, because cellular abnormalities are not restricted to one particular HLA phenotype, and targeted therapy requires some knowledge of the phenotype of the abnormal cells at issue.

The enzyme tyrosinase catalyzes the reaction converting

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tyrosine to dehydroxyphenylalanine or "DOPA" and appears to be expressed selectively in melanocytes (Muller et al., EMBOJ 7: 2715 (1988)). An early report of cDNA for the human enzyme is found in Kwon, U.S. Patent No. 4,898,814. A later report by Bouchard et al., J. Exp. Med. 169: 2029 (1989) presents a slightly different sequence. A great deal of effort has gone into identifying inhibitors for this enzyme, as it has been implicated in pigmentation diseases. Some examples of this literature include Jinbow, WO9116302; Mishima et al., U.S. Patent No. 5,077,059, and Nazzaropor, U.S. Patent 4,818,768. The artisan will be familiar with other references which teach similar materials.

U.S. Patent Application 08/081,673, filed June 23, 1993 and incorporated by reference, teaches that tyrosinase may be treated in a manner similar to a foreign antigen or a TRAP molecule - i.e., it was found that in certain cellular abnormalities, such as melanoma, tyrosinase is processed and a peptide derived therefrom forms a complex with HLA molecules on certain abnormal cells. These complexes were found to be recognized by cytolytic T cells ("CTLs"), which then lyse the The ramifications of this surprising and presenting cells. unexpected phenomenon were discussed. Additional peptides have now been found which also act as tumor rejection antigens presented by HLA-A2 molecules. These are described in Serial No. 08/203,054, filed February 28, 1994 and incorporated by reference.

It has now been found that additional peptides derived from tyrosinase are tumor rejection antigens in that they are presetend by MHC molecule HLA-B44, and are lysed by cytolytic T cells.

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# 5 BRIEF DESCRIPTION OF THE FIGURES

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Figure 1 describes, collectively, cell lysis studies. In particular:

Figure 1A shows lysis of cell line LB24-MEL;

Figure 1B shows lysis of cell line SK29-MEL;

Figure 1C shows lysis of cell line LB4.MEL;

Figure 1D shows lysis of cell line SK23.MEL;

Figure 1E shows lysis of cell line LE516.MEL;

Figure 1F shows lysis of cell line SK29-MEL.1.22 which has lost HLA-A2 expression;

Figure 1G shows lack of lysis of MZ2-MEL;

Figure 1H shows lysis studies on NK target K562;

Figure 1I shows lysis of the loss variant in Figure 1F after transfection with a gene for HLA-A2;

Figure 1J shows lysis of autologous EBV transformed B cells from patient SK29.

Figure 2 presents studies of TNF release of CTL IVSB.

Figure 3 depicts studies of TNF release of CTL 210/9.

Figure 4 depicts the recognition of the peptide YMNGTMSQV by cytolytic T cell clone CTL-IVSB but not cytolytic T cell clone CTL 2/9.

Figure 5 shows that the peptide YMNGTMSQV is not recognized by cytolytic T cell clone CTL 210/9.

Figure 6 shows the results obtained when TNF release assays were carried out on various cells, including those which present HLA-B44 on their surface.

Figure 7 shows, collectively, a series of chromium release assays using peptides described in this application on three different cell lines.

Figure 7A presents experiments where the peptide of SEQ ID NO: 4 was used.

Figure 7B shows results where the peptide of SEQ ID NO:  $5\ \text{was}$  used.

Figure 7C sets forth results obtained using SEQ ID NO: 2. In Figure 7, the symbol "O" is used for cell line T2, "O" for MZ2-MEL not presenting HLA-A2, and "O" for MZ2-MEL which has been transfected to present HLA-A2. Example 12

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5 elaborates on these tests.

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Figures 8A and 8B show work using a cell line which presents MHC molecule HLA-B44, and cytolytic T cell clone 22/31 ("CTL 22/31" hereafter). In figure 8A, the cell line ("Rosi EBV") was preincubated with monoclonal antibody W6/32, whereas in figure 8B, there was no preincubation.

# DETAILED DESCRIPTION OF PREFERRED EMBODIMENTS Example 1

Melanoma cell lines SK 29-MEL (also referred to in the literature as SK MEL-29) and LB24-MEL, which have been available to researchers for many years, were used in the following experiments.

Samples containing mononuclear blood cells were taken from patients SK29 (AV) and LB2 (these patients were also the source of SK 29-MEL and LB24-MEL, respectively). The melanoma cell lines were contacted to the mononuclear blood cell containing samples. The mixtures were observed for lysis of the melanoma cell lines, this lysis indicating that cytolytic T cells ("CTLs") specific for a complex of peptide and HLA molecule presented by the melanoma cells were present in the sample.

The lysis assay employed was a chromium release assay following Herin et al., Int. J. Cancer 39:390-396 (1987), the disclosure of which is incorporated by reference. The assay, however, is described herein. The target melanoma cells were grown in vitro, and then resuspended at 10° cells/ml in DMEM, supplemented with 10 mM HEPES and 30% FCS, and incubated for 45 minutes at 37°C with 200  $\mu$ Ci/ml of Na(51Cr)O<sub>4</sub>. cells were washed three times with DMEM, supplemented with 10 These were then resuspended in DMEM supplemented with 10 mM Hepes and 10% FCS, after which 100 ul aliquots containing 10³ cells, were distributed into microplates. Samples of PBLs were added in 100 ul of the same medium, and assays were carried out in duplicate. Plates were centrifuged for 4 minutes at 100g, and incubated for four hours at 37°C in a 5.5% of CO2 atmosphere.

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Plates were centrifuged again, and 100 ul aliquots of supernatant were collected and counted. Percentage of 51Cr release was calculated as follows:

%  $^{51}$ Cr release = <u>(ER-SR)</u> x 100 (MR-SR)

where ER is observed, experimental 51Cr release, SR is spontaneous release measured by incubating 103 labeled cells in 200 ul of medium alone, and MR is maximum release, obtained by adding 100 ul 0.3% Triton X-100 to target cells.

Those mononuclear blood samples which showed high CTL activity were expanded and cloned via limiting dilution, and were screened again, using the same methodology.

The same method was used to test target K562 cells. When EBV-transformed B cells (EBV-B cells) were used, the only change was the replacement of DMEM medium by Hank's medium, supplemented with 5% FCS.

These experiments led to isolation of CTL clone "IVSB" from patient SK29 (AV) and CTL clone 210/9 from patient LB24.

Figure 1 presents the results of these assays, in panels A, B, G and I. Specifically, it will be seen that both CTLs lysed both melanoma cell lines, and that there was no lysis of the K562 and EBVB cell lines.

#### Example 2

The CTLs described were tested against other melanoma cell lines to determine whether their target was shared by other melanoma cell lines. Lysis as described in Example 1 was studied for lines LB4.MEL, SK23.MEL (also known as SK MEL-23), and LE516.MEL. Figure 1, panels C, D and E shows that the clones did lyse these lines.

The tested lines are known to be of type HLA-A2, and the results suggested that the CTLs are specific for a complex of peptide and HLA-A2. This suggestion was verified by testing a variant of SK 29-MEL which has lost HLA-A2 expression. Figure 1, panel F shows these results. Neither clone lysed the HLA-loss variant. When the variant was transfected with the HLA-A2 gene of SK29-MEL, however, and retested, lysis was observed. Thus, it can be concluded that the presenting

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molecule is HLA-A2.

#### Example 3

Once the presenting HLA molecule was identified, studies were carried out to identify the molecule, referred to hereafter as the "tumor rejection antigen precursor" or "TRAP" molecule which was the source of the presented peptide.

To do this, total RNA was isolated from cell line SK29-MEL.1, which is a subclone of SK29-MEL. The RNA was isolated using an oligo-dT binding kit, following well recognized techniques. Once the total RNA was secured, it was transcribed into cDNA, again using standard methodologies. The cDNA was then ligated to EcoRI adaptors and cloned into the EcoRI site of plasmid pcDNA-I/Amp, in accordance with manufacturer's instructions. The recombinant plasmids were then electroporated into JM101  $\underline{E}$ .  $\underline{coli}$  (electroporation conditions: 1 pulse at 25  $\mu$ farads, 2500 V).

The transfected bacteria were selected with ampicillin  $(50~\mu\text{g/ml})$ , and then divided into 700 pools of 200 clones each. Each pool represented about 100 different cDNAs, as analysis showed that about 50% of plasmids contained an insert. Each pool was amplified to saturation, and plasmid DNA was isolated via alkaline lysis, potassium acetate precipitation and phenol extraction, following Maniatis et al., in Molecular Cloning: A Laboratory Manual (Cold Spring Harbor, N.Y., 1982). Cesium gradient centrifugation was not used.

### Example 4

The amplified plasmids were then transfected into eukaryotic cells. Samples of COS-7 cells were seeded, at 15,000 cells/well into tissue culture flat bottom microwells, in Dulbeco's modified Eagles Medium ("DMEM") supplemented with 10% fetal calf serum. The cells were incubated overnight at 37°C, medium was removed and then replaced by 30  $\mu$ l/well of DMEM medium containing 10% Nu serum, 400  $\mu$ g/ml DEAE-dextran, 100  $\mu$ M chloroquine, 100 ng of plasmid pcDNA-I/Amp-A2 and 100 ng of DNA of a pool of the cDNA library described supra. Plasmid pcDNA-I/Amp-A2 contains the HLA-A2 gene from SK29-MEL.

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Following four hours of incubation at 37°C, the medium was removed, and replaced by 50  $\mu$ l of PBS containing 10% DMSO. This medium was removed after two minutes and replaced by 200  $\mu$ l of DMEM supplemented with 10% of FCS.

Following this change in medium, COS cells were incubated for 48 hours at 37°C. Medium was then discarded, and 2000 cells of either of the described CTL clones were added, in 100  $\mu$ l of Iscove medium containing 10% pooled human serum. When clone 210/9 was used, the medium was supplemented with 25 U/ml of IL-2. Supernatant was removed after 24 hours, and TNF content was determined in an assay on WEHI cells, as described by Traversari et al., Immunogenetics 35: 145-152 (1992), the disclosure of which is incorporated by reference.

Of 700 wells tested with IVSB, 696 showed between 0.6 and 4 pg of TNF per ml. The remaining four wells contained between 10 and 20 pg/ml of TNF. Homologous wells tested with CTL 210/9 showed similar, clearly higher values. Figures 2 and 3 present these data.

# Example 5

Three of the four pools identified as high producers (numbers "123", "181" and "384") were selected for further experiments. Specifically, the bacteria were cloned, and 570 bacteria were tested from each pool. Plasmid DNA was extracted therefrom, transfected into a new sample of COS cells in the same manner as described supra, and the cells were again tested for stimulation of CTL 210/9 and CTL IVSB. A positive clone was found in pool 123 ("pl23.B2"), and one was found in pool 384 ("p384.C6"). Convincing evidence that the transfected cells were recognized by CTLs was obtained by carrying out a comparative test of COS cells transfected with cDNA and the HLA-A2 gene, and COS cells transfected only with TNF release in CTL supernatant was measured by HLA-A2. testing it on WEHI cells. The optical density of the surviving WEHI cells was measured using MTT. Results are presented in Table 1:

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#### Table 1

		CDNA (123.B2) + HLA-A2 DNA	no cDNA+ HLA-A2
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	Run 1	0.087	0.502
	Run 2	0.108	0.562

The values for WEHI OD's correspond to 24 pg/ml of TNF for cDNA and HLA-A2, versus 2.3 pg/ml for the control.

The plasmids from the positive clones were removed, and sequenced following art known techniques. A sequence search revealed that the plasmid insert was nearly identical to the cDNA for human tyrosinase, as described by Bouchard et al., J. Exp. Med. 169: 2029 (1989), the disclosure of which is incorporated by reference. Thus, a normally occurring molecule (i.e., tyrosinase), may act as a tumor rejection antigen precursor and be processed to form a peptide tumor rejection antigen which is presented on the surface of a cell, in combination with HLA-A2, thereby stimulating lysis by CTL clones. The nucleic sequence of the identified molecule is presented as SEQ ID NO: 1.

#### Example 6

Prior work reported by Chomez et al., Immunogenetics 35: 241 (1992) has shown that small gene fragments which contain a sequence coding for an antigenic peptide resulted in expression of that peptide. This work, which is incorporated by reference in its entirety, suggested the cloning of small portions of the human tyrosinase cDNA described supra and in SEQ ID NO: 1. Using the methodologies described in examples 1-5, various fragments of the cDNA were cotransfected with a gene for HLA-A2 in COS-7 cells, and TNF release assays were performed. These experiments led to identification of an approximately 400 base pair fragment which, when used in cotransfection experiments, provoked TNF release from

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cytolytic T cell clone CTL IVSB discussed supra, shown to be specific for HLA-A2 presenting cells. The 400 base fragment used corresponded to bases 711 to 1152 of SEQ ID NO: 1. amino acid sequence for which the fragment codes was deduced, and this sequence was then compared to the information provided by Hunt et al., Science 255: 1261 (1992), and Falk et al., Nature 351: 290 (1991), the disclosures of which are both incorporated by reference in their entirety. These references discuss consensus sequences for HLA-A2 presented peptides. Specifically, Hunt discusses nonapeptides, where either Leu or Ile is always found at the second position, Leu being the "dominant residue". The ninth residue is described as always being a residue with an aliphatic hydrocarbon side chain. Val is the dominant residue at this position. Hunt, discusses a strong signal for Leu and an intermediate signal for Met at the second position, one of Val, Leu, Ile or Thr at position 6, and Val or Leu at position 9, with Val being particularly On the basis of the comparison, nonapeptides were synthesized and then tested to see if they could sensitize HLA-A2 presenting cells. To do so, tyrosinase loss variant cell lines SK29-MEL 1.218 and T202LB were used. concentrations of the tested peptides were added to the cell lines, together with either of cytolytic T cell clone CTL IVSB or cytolytic T cell clone CTL 2/9. Prior work, described supra, had established that the former clone lysed tyrosinase expressing cells which present HLA-A2, and that the latter did not.

The tyrosinase loss variants were incubated for one hour in a solution containing  $^{51}$ Cr, at 37°C, either with or without anti HLA-A2 antibody MA2.1, which was used to stabilize empty HLA-A2 molecules. In the tests, cells were washed four times, and then incubated with varying dilutions of the peptides, from 100  $\mu$ M down to 0.01  $\mu$ M. After 30 minutes, effector cells were added at an E/T ratio of 40/1 and four hours later, 100 $\lambda$  of supernatant were collected and radioactivity counted.

Figure 4 shows the results obtained with nonapeptide

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5 Tyr Met Asn Gly Thr Met Ser Gln Val.

(SEQ ID NO: 2).

This peptide, referred to hereafter as SEQ ID NO: 2, corresponds to residues 1129-1155 of the cDNA sequence for tyrosinase presented in SEQ ID NO: 1. Complexes of HLA-A2 and this peptide are recognized by CTL clone IVSB.

In a parallel experiment, it was shown that CTL clone CTL 210/9, derived from patient LB24, did not recognize the complexes of HLA-A2 and the peptide of SEQ ID NO: 2, although it did recognize complexes of HLA-A2 and a tyrosinase derived peptide. Thus, tyrosinase is processed to at least one additional peptide which, when presented by HLA-A2 molecules, is recognized by CTL clones.

#### Example 7

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In a follow-up experiment, a second gene fragment which did not encode the peptide of SEQ ID NO: 2 was used. This fragment began at base 1 and ended at base 1101 of SEQ ID NO: 1 (i.e. the EcoRI-SphI fragment). Cytolytic T cell clone CTL 210/9, discussed <u>supra</u>, was tested against COS-7 cells transfected with this fragment in the manner described <u>supra</u>. CTL IVSB was also tested. These results, showed that LB24-CTL 210/9 recognized an antigen on the surface of HLA-A2 expressing cells transfected with this fragment, but CTL IVSB did not. Thus, a second tumor rejection antigen peptide is derived from tyrosinase.

#### Example 8

In order to further define the tumor rejection antigen recognized by LB24-CTL 210/9, the following experiments were carried out.

A second fragment, corresponding to bases 451-1158 of SEQ ID NO: 1 was transfected into COS cells together with a gene for HLA-A2, and TNF release assays were carried out. This sequence provoked TNF release from clone SK29-CTL IVSB (20 pg/ml), but not from LB24-CTL 210/9 (3.8 pg/ml). These results confirmed that the two CTL clones recognize different peptides, and that the peptide recognized by LB24-CTL 210/9 must be encoded by region 1-451.

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# 5 Example 9

The tyrosinase derived peptide coded for by cDNA fragment 1-451 was analyzed for consensus sequences known to bind HLA-A2. The peptides corresponding to these consensus sequences were synthesized, and tested for their ability to sensitize HLA-A2 presenting cells. To do so, two tyrosinase negative melanoma cell lines were used (i.e., NA8-MEL, and MZ2-MEL 2.2 transfected with HLA-A2), and cell line T2, as described by Salter et al, Immunogenetics 21: 235-246 (1985)).

The cells were incubated with <sup>51</sup>Cr, and monoclonal antibody MA.2.1, which is specific for HLA-A2 for 50 minutes at 37°C, followed by washing (see Bodmer et al., Nature 342: 443-446 (1989), the disclosure of which is incorporated by reference in its entirety). Target cells were incubated with various concentrations of the peptides, and with either of LB 24-CTL clones 210/5 or 210/9. The percent of chromium release was measured after four hours of incubation.

The peptide Met Leu Leu Ala Val Leu Tyr Cys Leu Leu (SEQ ID NO: 3) was found to be active.

In further experiments summarized here, CTL-IVSB previously shown to recognize YMNGTMSQV, did not recognize the peptide of SEQ ID NO: 3.

The results are summarized in Tables 2-4 which follow:

#### Table 2

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			Peptide	
		YMNGTMSQV		MLLAVLYCLL
		(1120-1155)		(25-54)
	SK29-CTL-IVSB	+		-
35	LB24-CTL-210/5	-		+
	LB24-CTL-210/9	_		+

## Table 3

3/93-Lysis of MZ2-2.2-A2 sensitized with tyrosinase peptides by LB24-CTL 210/5 and 210/9, and SK29-CTL IVSB

Effectors	Peptides	Dose	MZ2.2.2-A2 + anti-A2*	
LB24-CTL 210/5 (44.1)	MLLAVLYCLL (LAUS	10μM 17-5)	18 3 1	17 16
·	YMNGTMSQV (MAINZ)	30M 10 3	1 1 1	
LB24-CTL 210/9 (30:1)	MLLAVLYCLL (LAUS		18 3 1	17 15
	YMNGTMSQV (MAINZ)	30M 10 3	1 1 1	
SK29-CTL IVSB (40:1)	MLLAVLYCLL (LAUS	10μM 17-5)	1 3 1	1 1
	YMNGTMSQV (MAINZ)	30μM 10 3	68 68 62	

<sup>\*</sup> Target cells were incubated with Cr51 and mono-Ab MA2.1 (anti-HLA-A2) for 50 min, then washed 3 times.

CTL cells were added at the indicated (E:T) ratio.
The % specific Cr51 release was measured after 4h incubation

They were incubated with various concentrations of peptides for 30 min

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Table 4

8j93 : Test of tyrosinase peptides recognized by LB24-CTL 210/5 and 210/9 or SK29-CTL IVSB

( % Cr51 specific release)

Effectors	Peptides	Dose	NA8-MEL •	MZ2-2.2: A2	Т2
LB24-CTL. 210/5 ( 41:1)	MLLAVLYCLL (LAUS 17-5)	10μM 3 1 300nM 100 30	30 23 17 6 2	31 27 20 17 8 5	36 35 26 16 5
	. 0		0	0	. 0
LB24-CTL. 210/9 ( 26:1 )	MLLAVLYCLL (LAUS 17-5)	10µM 3 1 300nM 100 30	14 13 9 3 1	19 17 14 9 1	21 20 13 5 1 0
	0		• 0	1	0
SK29-CTL. IVSB ( 42:1 )	YMNGTMSQV (MAINZ)	10μM 3 1 300nM 100 30 10 3	46 38 27 14 3 1 0	46 44 40 22 13 9 3	59 52 46 34 21 10 3
	. 0		0	4	0
spt. rel. max-spt %	* • .		339 2694 11	259 1693 13	198 1206 14

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# 5 Example 10

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Additional experiments were carried out using CTL clone 22/31. This clone had previously been shown to lyse subline MZ2-MEL.43 from autologous melanoma cell line MZ2-MEL, but did not lyse other sublines, such as MZ2-MEL 3.0 and MZ2-MEL 61.2, nor did it lyse autologous EBV transformed B cells, or killer cell line K562 (see Van den Eynde et al., Int. J. Cancer 44: 634-640 (1989)). The antigen presented by MZ2-MEL.43 is referred to as antigen C.

In prior work including that reported in the parent of this application, it was found that the tyrosinase gene encodes an antigen recognized by autologous CTLs on most HLA-A2 expressing melanomas. Expression of this gene in sublines of cell line MZ2-MEL was tested by PCR amplification. Clone MZ2-MEL.43 was found to be positive, whereas other MZ2-MEL clones, such as MZ2-MEL.3.0 were negative. Correlation of expression of the tyrosinase gene, and antigen MZ2-C, suggested that MZ2-C might be a tumor rejection antigen derived from tyrosinase, and presented by an HLA molecule expressed by MZ2-MEL. This cell line does not express HLA-A2, which would indicate that if a tyrosinase derived peptide were presented as a TRA, a second HLA molecule was implicated.

Studies were carried out to identify which HLA molecule presented antigen C to CTL 22/31. To determine this, cDNA clones of the HLA molecules known to be on the cell surface, i.e., HLA-A29, HLA-B37, HLA-B 44.02, and HLA-C clone 10, were isolated from an MZ2-MEL.43 cDNA library, and then cloned into expression vector pcDNAI/Amp. Recipient COS 7 cells were then transfected with one of these constructs or a construct containing HLA-A1, plus cDNA coding for tyrosinase (SEQ ID NO: 1). The contransfection followed the method set forth above. One day later CTL 22/31 was added, and 24 hours later, TNF release was measured by testing cytotoxicity on WEHI-164-13, following Traversari et al, supra. Figure 6 shows that TNF was released by CTL 22/31 only in the presence of cells transfected with both HLA-B44 and tyrosinase. The conclusion to be drawn from this is that HLA-B44 presents a tyrosinase

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5 derived tumor rejection antigen.

### Example 11

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The experiments described supra showed, inter alia, that the decamer MLLAVLYCLL effectively induced lysis of HLA-A2 presenting cells. It is fairly well accepted that MHC molecules present nonapeptides. To that end, experiments were carried out wherein two nonamers were tested, which were based upon the decapeptide which did give positive results. Specifically, either the first or tenth amino acid was omitted to create two peptides, i.e.:

Met Leu Leu Ala Val Leu Tyr Cys Leu

(SEQ ID NO: 4)

Leu Leu Ala Val Leu Tyr Cys Leu Leu

(SEQ ID NO: 5).

These peptides were tested in the same way the decapeptide was tested, as set forth in the prior examples at concentrations ranging from 10  $\mu\text{M}$  to 1 nM. Three presenting cells were used. As summarized in Table 5, which follows, "T2" is a mutant human cell line, "CEMX721.174T2" as described by Salter, Immunogenetics 21: 235(1985). This line presents HLA-A2. "G2.2" is a variant of the cell line MZ2-MEL. The variant has been transfected with a gene coding for HLA-A2. The abbreviation "G2.2.5" stands for a variant which does not express HLA-A2. All cells were incubated with monoclonal antibody MA2.1 prior to contact with the cytolytic T cell This procedure stabilizes so-called "empty" MHC molecules, although the mechanism by which this occurs is not well understood and effector CTLs 210/5 and 210/9 were both The results are set forth in Table 5, which follows. They show that at a concentration of 10  $\mu$ M, the nonamer of SEQ ID NO: 4 was twice as effective when used with CTL clone 210/5, and four times as effective with clone 210/9 whereas the nonamer of SEQ ID NO: 5 was ineffective at inducing lysis. Example 12

In further experiments, chromium release assays were carried out using the peptides of SEQ ID NOS: 4 and 5, as well as SEQ ID NO: 2. The target cells were allogeneic melanoma

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cells, i.e., MZ2-MEL, previously transfected with HLA-A2, and cell line T2, which presents HLA-A2, but has an antigen processing defect which results in an increased capacity to present exogenous peptides (Cerundolo et al., Nature 345: 449 (1990)). All cells were pretriated with monoclonal antibody MA2.1 for fifty minutes. The cells were incubated with the peptide of choice, for 30 minutes, at various concentrations. Then, one of CTL clones 210/9 and ISVB was added in an effector: target ratio of 60. Chromium release was measured after four hours, in the manner described supra.

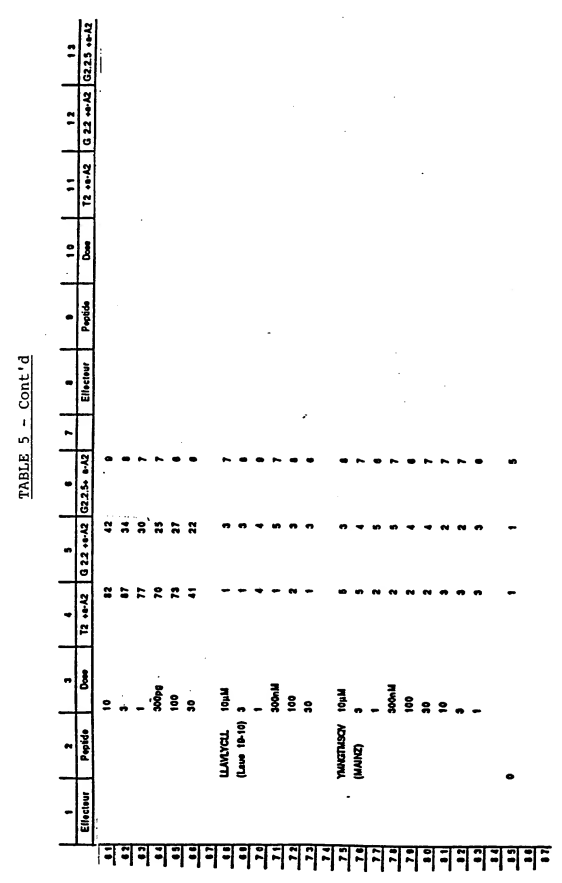
The results are presented in figure 7, i.e., figures 7A-7C. The peptide of SEQ ID NO: 4 sensitized cells to CTL 210/9, while SEQ ID NO: 5 did not. SEQ ID NO: 6 sensitized cells to CTL IVSB, as already noted in previous examples.

-	~	-	•	5	•	7 8	•	10	11	12	=
Effecteur	Pepide	Dose	T2 +e-A2	Q 2.2 +4-A2	G2.2.5+ 4-A2	Elfecteur	Peptide	Dose	12 .0.42	G 22 +4-A2	G2.2.5 ++- A2
1 DAGI 210/5	METANTAL	10µM	80	32		SK29 IVSB	MITANTAIT	10401		6	
2 50:1	(LAUS 17-6)	•	7	32	•	1:09	(LAUS 17-5)		•	~	
6		-	30	28	•			-	a	•	
=		300nM	33	=	•			3000M	•	-	
<b>S</b>		100	54	•	•			100	-	~	
		90	13	•	•			90	-	•	
-		•	•	<b>5</b>				10	.0	•	
		•	~	~ ~	•			•	~	•	
			~	-	•			_	~	•	
ि		300bg	-	•	•			300pg	-	~	
<del>≠</del>  ≈	MILAMYCL	10pM	•	8	^		MILANTYCL	10nM	~	•	
<u> </u>	(LAUS 18-5)	•	11	9	•	. <b>.</b>	(LAUS 18-5)			•	
-			60	8	10			-	•	•	
1 5		300nM	80	9	n			300nM	-	•	
٥		9	5	26	•			100	-	•	
<b>—</b>		90	11	;	•		-	30	•	•	
=		2			•			0	•	•	
		•	7.		<b>~</b>			•	-	•	
20		_	7.8		***			_	-	•	
~		300pg	7		₩7			300pg	•	n	
2 2	LAMYCLL	10µM	•	-	•		<b>LLAMYCIL</b>	N <sub>4</sub> 01	•	•	
2.4	(Laue 19-10)	•	•	8	•		(Laue 19-10)	•	•	•	
2.5		-	•	•	•			-	-	•	
2 6		300nM	•	8	<b>▼</b>			300m	Ŋ	•	
22		00	-	~	•			00	-	₩7	
20		30	-	-	~			30	-	◄	
9 0	YMNGTMSOV	. Myot	•	•	₹ <b>→</b>		YMAGTASQV	Nugot	7.8	8	

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TABLE 5 - Cont'd

1		~	- -	-	·	•	~	•	•	•	-	12	2
MALANYTEL 10pH   1	Effecteur		Dose	T2 +6-A2	G 2.2 +6-A2	G2.2.5+ e-A2	$\prod$	Effecteur	Peptide	Dose	T2 +e-A2	G 22 **- 1/2	G2.25 ++- A2
1		(MANZ)	•	•	-				(MAINZ)		73		-
100   1   2   2   100   100   4   100			-	~	~	•				-	93		1
100   0   1   6   100   25   100			300nM	-	•	~				300nM	\$	•	•
10   2   4   30   22   18   19   10   23   18   19   10   23   18   19   10   23   18   19   19   19   19   19   19   19			901	•	-	•				100	4	7	,
10   0   0   10   10   10   10   10			90	•	~	•				30	30	•	~
1			2	•		**				01	23	=	•
1			•	•		•				•	=	=	•
100			_	~		46				-	•	9	•
NALIANIYCLI   194M   28   23   8   20   2   7   20   2   7   20   2   2   2   2   2   2   2   2										300pg	•	^	•
MALANYCAL   Oph M		•		•	•	^				100	•	1	~
MALANLYCK   1944   20   23   6   0   2   3   1   1   1   1   1   1   1   1   1										30	~	7	•
LAUS 17-5  3	AQ1 210/0		10µM	20	23	•	. •						
1	::		•	20	23	•	<i>-</i>			•	N	•	•
300nM         13         16         441           100         10         9         9         15         1				•	22	a							
100         10         6         max-opt         1522           30         6         7         %         15			300m	13	=	•	-	opt ref.			:		105
30     6     7     %       10     3     4     6       3     6     6       10     7     3     6       300pg     1     4     7       100     1     4     7       30     1     4     7       10µM     82     75     10       100     82     75     10       100     87     6     6       100     87     6       100     83     53     5			001	9	•	•	_	mex-ept			1033		1686
10 30 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5			90	•	•	_		*					2
3 1 100 100 100 10µM 3 1 1000 100 100 100 100 100 100 100 10			2	•	~	•						•	
100pg 1 100 1 10µM 96 1 89 1 60 87			•	**	•	<b>50</b>							
300pg 100 10µM 98 1 100nM 95 100 87			_	1	•	•							
100 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1			300pg	-	•	•							
10µM 96 1 1 90 1 1 90 1 1 90 1 91 91 91 91 91 91 91 91 91 91 91 91 9			100	•	•	•							
10µM 88 1 89 1000 87 30 83			30	-	•	^		٠					
3 1 300nM 85 100 87		MILANYEL	10µM	•	13	12							
0 0 0		(LAUS 16-5)	•	93	75	0					-		
8 <b>9</b> 8			_	•	7.								
F 60			300nM	50	01	•					•		
50			, 001	11	63	•							
		·	. 00	60	53	<b>10</b>							



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#### Example 13

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Work which followed up on the experiments set forth in example 10 was then carried out, in an effort to define the antigenic peptide presented by HLA-B44. To do so, cDNA sequences corresponding to fragments of the tyrosinase cDNA sequence were cotransfected, together with a gene coding for HLA-B44, into COS-7 cells. The protocol is essentially that described in example 6, supra. The cytolytic T cell clone 22/31, discussed <u>supra</u>, was used. TNF release was determined. Two fragments, i.e., base fragments 1-611, and 427-1134 induced TNF release. This suggested that the presented peptide was in the overlapping region. As a result of this observation, shorter fragments were tested. Fragments beginning at positions 385, 442, 514 and 574 were able to induce TNF release, while fragments starting at positions 579 and 585 were not. These observations, in turn, suggested the synthesis, following standard methodologies, of a 13 amino acid peptide beginning at position 574.

This peptide was then used in experiments to determine whether it induced lysis by CTL 22/31. Table 6, which follows, shows that the 13-mer rendered two EBV transfect d cell lines which express HLA-B44 sensitive to lysis.

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# Table 6

10F94-tyros 1	IS-mer eur EBV-I			
. 1	1 2	3	4	
Eff	ector Dose pept 13 AA	Rosi -EBV	MZ2 - EBV	
1 MZ2-C	TL-22/31 SEWRDIDFAHEA			
3 60:1	<b>3</b> 0µМ	83	71	
4	10	85	72	
	3	77	66	
	1	79	63	
7	<b>3</b> 00nM	60	33	
	100	44	17	
9	30	21	• 4	
1 0	10	9	5	•
11	3	10	6	
12	0	10	6	
1 4 1 5 spt.re	ıi.	393	472	
1 6 max.r	el.	1898	1792	
17%		23	26	

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As a follow up, even shorter peptides were tested. A decamer corresponding to nucleotide bases 574-604, i.e.

Ser Glu Ile Trp Arg Asp Ile Asp Phe Ala

(SEQ ID NO: 7)

did provoke lysis, as did peptide:

Ser Glu Ile Trp Arg Asp Ile Asp Phe

(SEQ ID NO: 8)

The nonamer:

Glu Ile Trp Arg Asp Ile Asp Phe Ala

(SEQ ID NO: 9)

in contrast, was not recognized. Table 7, which follows,

summarizes these results, which are also depicted in figur 8.

The only other peptide reported to be bound by HLA-B44 is

Glu Glu Asn Leu Leu Asp Phe Val Arg Phe

(SEQ ID NO: 10)

as reported by Burrows et al., J. Virol 64: 3974 (1990). The data described supra suggest that Glu at second position and Phe in ninth position may represent anchor residues for HLA-B44.

Table 7

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Dose 1µM 300nM	+W SEIWRDIDFA 91 76	+W SEIWRDIDF 93 81
	76	81
30011141		
100	43	73
	17	37
	. 4	12
	3	4
3	2	. 4
1		1
	100 30 10 3 1	30 17 10 4 3 3 1

24

TABLE 7 - Cont'd

1 8	6	7	8
+W EIWRDIDFA	-W SEIWRDIDFA	-W SEIWRDIDF	-W EIWRDIDFA
7	98	99	11
4	77	074	<del></del>
2	45	64	. 8
0	15	21	. 6
1	5	8	4
1	0	7	. 2
		3	
		. 2	

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The foregoing experiments demonstrate that tyrosinase is processed as a tumor rejection antigen precursor, leading to formation of complexes of the resulting tumor rejection antigens with a molecule on at least some abnormal cells, for example, melanoma cells with HLA-A2 or HLA-B44 phenotype. The complex can be recognized by CTLs, and the presenting cell This observation has therapeutic and diagnostic ramifications which are features of the invention. respect to therapies, the observation that CTLs which are specific for abnormal cells presenting the aforementioned produced, suggests various therapeutic complexes are One such approach is the administration of CTLs approaches. specific to the complex to a subject with abnormal cells of the phenotype at issue. It is within the skill of the artisan to develop such CTLs in vitro. Specifically, a sample of cells, such as blood cells, are contacted to a cell presenting the complex and are capable of provoking a specific CTL to proliferate. The target cell can be a transfectant, such as a COS cell of the type described supra. These transfectants present the desired complex on their surface and, when combined with a CTL of interest, stimulate its proliferation. So as to enable the artisan to produce these CTLs, vectors containing the genes of interest, i.e., pcDNA-1/Ampl (HLA-A2), (human tyrosinase), have been deposited in accordance with the Budapest Treaty at the Institut Pasteur, under Accession Numbers I1275 and I1276, respectively. cells, such as those used herein are widely available, as are other suitable host cells.

To detail the therapeutic methodology, referred to as adoptive transfer (Greenberg, J. Immunol. 136(5): 1917 (1986); Reddel et al., Science 257: 238 (7-10-92); Lynch et al., Eur. J. Immunol. 21: 1403-1410 (1991); Kast et al., Cell 59: 603-614 (11-17-89)), cells presenting the desired complex are combined with CTLs leading to proliferation of the CTLs specific ther to. The proliferated CTLs are then administered to a subject with a cellular abnormality which is characterized by certain of the abnormal cells presenting the

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particular complex. The CTLs then lyse the abnormal cells, thereby achieving the desired therapeutic goal.

The foregoing therapy assumes that at least some of the cells present one or more of subject's abnormal HLA/tyrosinase derived peptide complexes. This can be determined very easily. For example CTLs are identified using the transfectants discussed supra, and once isolated, can be used with a sample of a subject's abnormal cells to determine If lysis is observed, then the use of lysis <u>in vitro</u>. specific CTLs in such a therapy may alleviate the condition associated with the abnormal cells. Α less methodology examines the abnormal cells for their HLA phenotype, using standard assays, and determines expression of tyrosinase via amplification using, e.g., PCR. The fact that a plurality of different HLA molecules present TRAs derived from tyrosinase increases the number of individuals who are suitable subjects for the therapies discussed herein.

Adoptive transfer is not the only form of therapy that is available in accordance with the invention. CTLs can also be provoked in vivo, using a number of approaches. One approach, i.e., the use of non-proliferative cells expressing the complex, has been elaborated upon supra. The cells used in this approach may be those that normally express the complex, such as irradiated melanoma cells or cells transfected with one or both of the genes necessary for presentation of the complex. Chen et al., Proc. Natl. Acad. Sci. USA 88: 110-114 (January, 1991) exemplifies this approach, showing the use of transfected cells expressing HPVE7 peptides in a therapeutic regime. Various cell types may be used. Similarly, vectors carrying one or both of the genes of interest may be used. Viral or bacterial vectors are especially preferred. In these systems, the gene of interest is carried by, e.g., a Vaccinia virus or the bacteria BCG, and the materials de facto "infect" The cells which result present the complex of host cells. interest, and are recognized by autologous CTLs, which then A similar effect can be achieved by combining proliferate. tyrosinase itself with an adjuvant to facilitate incorporation

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into HLA-A2 presenting cells. The enzyme is then processed to yield the peptide partner of the HLA molecule.

The foregoing discussion refers to "abnormal cells" and "cellular abnormalities". These terms are employed in their broadest interpretation, and refer to any situation where the cells in question exhibit at least one property which indicates that they differ from normal cells of their specific type. Examples of abnormal properties include morphological and biochemical changes, e.g. Cellular abnormalities include tumors, such as melanoma, autoimmune disorders, and so forth.

The invention also provides a method for identifying precursors to CTL targets. These precursors are referred to as tumor rejection antigens when the target cells are tumors, but it must be pointed out that when the cell characterized by abnormality is not a tumor, it would be somewhat misleading to refer to the molecule as a tumor rejection antigen. Essentially, the method involves identifying a cell which is the target of a cytolytic T cell of the type discussed supra. Once such a cell is identified, total RNA is converted to a cDNA library, which is then transfected into a cell sample capable of presenting an antigen which forms a complex with a relevant HLA molecule. The transfectants are contacted with the CTL discussed supra, and again, targeting by the CTL is observed (lysis and/or TNF production). These transfectants which are lysed are then treated to have the cDNA removed and sequenced, and in this manner a precursor for an abnormal condition, such as a tumor rejection antigen precursor, can be identified.

Other aspects of the invention will be clear to the skilled artisan and need not be repeated here.

The terms and expressions which have been employed are used as terms of description and not of limitation, and there is no intention in the use of such terms and expressions of excluding any equivalents of the features shown and described or portions thereof, it being recognized that various modifications are possible within the scope of the invention.

- 1) GENERAL INFORMATION:
  - (i) APPLICANTS: Wölfel, Thomas; Van Pel, Aline; Brichard, Vincent; Boon-Falleur, Thierry
  - (ii) TITLE OF INVENTION: ISOLATED, TYROSINASE DERIVED PEPTIDES AND USES THEREOF
  - (iii) NUMBER OF SEQUENCES: 10
  - (iv) CORRESPONDENCE ADDRESS:
    - (A) ADDRESSEE: Felfe & Lynch
    - (B) STREET: 805 Third Avenue
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    - (D) STATE: New York
    - (E) COUNTRY: USA
    - (F) ZIP: 10022
  - (v) COMPUTER READABLE FORM:
    - (A) MEDIUM TYPE: Diskette, 5.25 inch, 360 kb storage
    - (B) COMPUTER: IBM PS/2
    - (C) OPERATING SYSTEM: PC-DOS
    - (D) SOFTWARE: Wordperfect
  - (vi) CURRENT APPLICATION DATA:
    - (A) APPLICATION NUMBER: 08/233,305
    - (B) FILING DATE: 26-APRIL-1994
    - (C) CLASSIFICATION: 514
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      - (A) APPLICATION NUMBER: 08/203,054
      - (B) FILING DATE: 28-FEB-1994
    - (vii) PRIOR APPLICATION DATA:
      - (A) APPLICATION NUMBER: 08/081,673
      - (B) FILING DATE: 23-JUNE-1993

- (vii) PRIOR APPLICATION DATA:
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  - (B) FILING DATE: 28-APRIL-1993
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  - (A) APPLICATION NUMBER: 07/994,928
  - (B) FILING DATE: 22-DEC-1992
- (viii) ATTORNEY/AGENT INFORMATION:
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(2)	INFORMATION FOR SEQ ID NO: 1:
	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 1894 base pairs
	(B) TYPE: nucleic acid
٠	(C) STRANDEDNESS: single
	(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

GGA AGA ATG CTC CTG GCT GTT TTG TAC TGC CTG CTG TGG AGT TTC CAG 48 Gly Arg Met Leu Leu Ala Val Leu Tyr Cys Leu Leu Trp Ser Phe Gln -15 -10 -5 ACC TCC GCT GGC CAT TTC CCT AGA GCC TGT GTC TCC TCT AAG AAC CTG 96 Thr Ser Ala Gly His Phe Pro Arg Ala Cys Val Ser Ser Lys Asn Leu 1 ATG GAG AAG GAA TGC TGT CCA CCG TGG AGC GGG GAC AGG AGT CCC TGT 144 Met Gly Lys Glu Cys Cys Pro Pro Trp Ser Gly Asp Arg Ser Pro Cys 15 20 GGC CAG CTT TCA GGC AGA GGT TCC TGT CAG AAT ATC CTT CTG TCC AAT 192 Gly Gln Leu Ser Gly Arg Gly Ser Cys Gln Asn Ile Leu Leu Ser Asn 35 40 45 GCA CCA CTT GGG CCT CAA TTT CCC TTC ACA GGG GTG GAT GAC CGG GAG 240 Ala Pro Leu Gly Pro Gln Phe Pro Phe Thr Gly Val Asp Asp Arg Glu 50 55 60 TCG TGG CCT TCC GTC TTT TAT AAT AGG ACC TGC CAG TGC TCT GGC AAC 288 Ser Trp Pro Ser Val Phe Tyr Asn Arg Thr Cys Gln Cys Ser Gly Asn 70 65 TTC ATG GGA TTC AAC TGT GGA AAC TGC AAG TTT GGC TTT TGG GGA CCA 336 Phe Met Gly Phe Asn Cys Gly Asn Cys Lys Phe Gly Phe Trp Gly Pro 80 85 90 AAC TGC ACA GAG AGA CGA CTC TTG GTG AGA AGA AAC ATC TTC GAT TTG 384 Asn Cys Thr Glu Arg Arg Leu Leu Val Arg Arg Asn Ile Phe Asp Leu 95 100 AGT GCC CCA GAG AAG GAC AAA TTT TTT GCC TAC CTC ACT TTA GCA AAG 432 Ser Ala Pro Glu Lys Asp Lys Phe Phe Ala Tyr Leu Thr Leu Ala Lys

120

125

115

CAT	ACC	ATC	AGC	TCA	GAC	TAT	GTC	ATC	CCC	ATA	GGG	ACC	TAT	GGC	CAA	480
His	Thr	Ile	Ser	Ser	Asp	Tyr	Va 1	Пe	Pro	Ile	Gly	Thr	Tyr	Gly	Gln	
			130					135					140			
ATG	AAA	AAT	GGA	TCA	ACA	CCC	ATG	TTT	AAC	GAC	ATC	AAT	ATT	TAT	GAC	528
Met	Lys	Aśn	Gly	Ser	Thr	Pro	Met	Phe	Asn	Asp	Пe	Asn	IJе	Tyr	Asp	
		145					150				-	155				
CTC	TTT	GTC	TGG	ATG	CAT	TAT	TAT	GTG	TCA	ATG	GAT	GCA	CTG	CTT	GGG	576
Leu	Phe	Va 1	Trp	Ile	His	Tyr	Tyr	Val	Ser	Met	Asp	Ala	Leu	Leu	Gly	
	160					165					170					
GGA	TCT	GAA	ATC	TGG	AGA	GAC	ATT	GAT	TTT	GCC	CAT	GAA	GCA	CCA	GCT	624
Gly	Tyr	Glu	Ile	Trp	Arg	Asp	Ilе	Asp	Phe	Ala	His	Glu	Ala	Pro	Ala	
175					180					185					190	
TTT	CTG	CCT	TGG	CAT	AGA	CTC	TTC	TTG	TTG	CGG	TGG	GAA	CAA	GAA	ATC	672
Phe	Leu	Pro	Trp	His	Arg	Leu	Phe	Leu	Leu	Arg	Trp	Glu	Gln	Gly	Ile	
				195					200					205		•
								TTC								720
Gln	Lys	Leu	Thr	Gly	Asp	Gly	Asn	Phe	Thr	I le	Pro	Tyr		Asp	Trp .	
			210					215	•				220			
								TGC								768
Arg	Asp		Glu	Lys	Cys	Asp		Cys	Thr	Asp	Gly		Met	Gly	Gly	
		225					230					235			<b>T</b> 00	016
-								CTC								816
Gln		Pro					Leu	Leu	Ser	Pro		Ser	Phe	Pne	Ser	
	240				<b>TOT</b>			TTO			250	886	6.00	CAT	CAC	064
								TTG								864
	irp	Gin	He	va i		5er	Arg	Leu	GIU		ıyr	ASII	Ser	піз	270	
255	<b>TT</b> 4	<b>T</b> 00		004	260	ccc	CAC	CCA	ССТ	265	ccc	ССТ	AAT	ССТ		912
								GGA								312
ser	Leu	Lys	ASTI		Inr	Pro	Giu	ч	280	Leu	AI 9	AI Y	ASII	285	Gly .	
	CAT	CAC		275	ΛĆΛ	۸۵۲	CCA	ACC		ccc	TCT	ΤCΔ	CCT		GTA	960
								Arg								300
ASTI	піз	ASÞ	290		AI 9	1111	710	295			361	JCI	300		74.	
CAA	TTT	TOO			TTC	Δ۲۲	CAA	TAT		TOT	GGT	TCC			ΑΑΑ	1008
								Tyr								
ulu	riie	305		261			310				,	315			-3-	
GCT	GCC			AGC	TTT	AGA	-		CTG	GAA	GGA			AGT	CCA	1056
							_									

Ala	Ala	Asn	Phe	Ser	Phe	Arg	Asn	Thr	Leu	Glu	Gly	Phe	Als	Ser	Pro	
	320					325					330					
			ATA													1104
Leu	Thr	Gly	Ile	Ala	Asp	Ala	Ser	G1n	Ser	Ser	Met	His	Asn	Ala	Leu	
335					340					345					350	
			ATG													1152
His	Ilе	Tyr	Met	Asn	Gly	Thr	Met	Ser	Gln	Met	Gln	Gly	Ser		Asn	
				355					360					365		
			TTC													1200
Asp	Pro	IЪе	Phe	Leu	Leu	His	His	Ala	Phe	Va 1	Asp	Ser	I је	Phe	Glu	
			370					375					380			
			CAA													1248
Gln	Trp	Leu	Arg	Arg	His	Arg		Leu	Gln	Glu	Val		Pro	Glu	Ala	
		385					390					395				
			ATT													1296
Asn		Pro	Ile	Gly	His		Arg	Glu	Ser	Tyr		Val	Pro	Phe	116	
	400					405					410				200	
			AGA													1344
Pro	Leu	Tyr	Arg	Asn		Asp	Phe	Phe	He			Lys	ASP	Leu		
415					420			704	040	425		TOT	***	C A A	430	1202
			AGC													1392
Tyr	Asp	Tyr	Ser		Leu	Gin	Asp	Ser			ASP	5er	Pne	445	ASP	
				435	<b>TT</b> 0		C A A	000	440		ATC	TCC	TCA		רדר	1440
															CTC	1440
ıyr	1 1e	Lys			Leu	ыу	GIII	455		mi y	1116	ПР	460		Leu	
CTT			450		CTA	ccc	ברר			ΔΓΤ	ecc	ств			GGG	1488
															Gly	
Leu	GIY	465		Met	Val	uiy	470		LCU	• • • • • • • • • • • • • • • • • • • •	,,,,	475			,	
стт	CTC			י רדם	TGT	. CET			AGA	AAG	CAG			GAA	GAA	1536
															Glu	
Leu	480		Leu	Leu	0,3	485		-,-	, ,,,	, -,-	490					
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Ser His Leu 513

TAAAAGGCTT	AGGCAATAGA	GTAGGGCCAA	AAAGCCTGAC	CTCACTCTAA	CTCAAAGTAA	1653
TGTCCAGGTT	CCCAGAGAAT	ATCTGCTGGT	ATTTTTCTGT	AAAGACCATT	TGCAAAATTG	1713
TAACCTAATA	CAAAGTGTAG	CCTTCTTCCA	ACTCAGGTAG	AACACACCTG	TCTTTGTCTT	1773
GCTGTTTTCA	CTCAGCCCTT	TTAACATTTT	CCCCTAAGCC	CATATGTCTA	AGGAAAGGAT	1833
GCTATTTGGT	AATGAGGAAC	TGTTATTTGT	ATGTGAATTA	AAGTGCTCTT	ATTTTAAAAA	1893
A	•					1894

- (2) INFORMATION FOR SEQ ID NO: 2:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 9 amino acid residues
    - (B) TYPE: amino acid
      - (D) TOPOLOGY: single
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

Tyr Met Asn Gly Thr Met Ser Gln Val
5

- (2) INFORMATION FOR SEQ ID NO: 3:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 10 amino acid residues
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: single
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

Met Leu Leu Ala Val Leu Tyr Cys Leu Leu 5 10

- (2) INFORMATION FOR SEQ ID NO: 4:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 9 amino acid residues
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

## Met Leu Leu Ala Val Leu Tyr Cys Leu

5

- (2) INFORMATION FOR SEQ ID NO: 5:
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    - (A) LENGTH: 9 amino acid residues
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:

Leu Leu Ala Val Leu Tyr Cys Leu Leu 5

- (2) INFORMATION FOR SEQ ID NO: 6:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13 amino acid residues
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:-

Ser Glu Ile Trp Arg Asp Ile Asp Phe Ala His Glu Ala 5 10

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- (2) INFORMATION FOR SEQ ID NO: 7:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 10 amino acid residues
    - (B) TYPE: amino acid
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  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:

Ser Glu Ile Trp Arg Asp Ile Asp Phe Ala

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- (2) INFORMATION FOR SEQ ID NO: 8:
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    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:

Ser Glu Ile Trp Arg Asp Ile Asp Phe

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- (2) INFORMATION FOR SEQ ID NO: 9:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 9 amino acid residues
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:

Glu Ile Trp Arg Asp Ile Asp Phe Ala

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- (2) INFORMATION FOR SEQ ID NO: 10:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 10 amino acid residues
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:

Glu Glu Asn Leu Leu Asp Phe Val Arg Phe

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#### We claim:

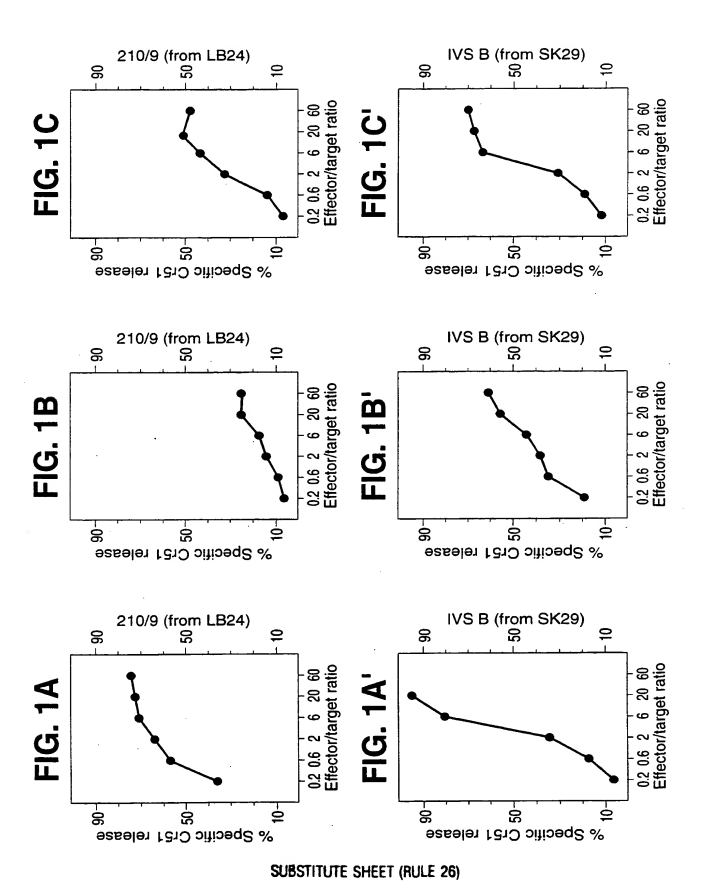
- 1. Method for identifying a candidate for treatment with a therapeutic agent specific for complexes of an MHC molecule and a tyrosinase derived peptide selected from the group consisting of SEQ ID NO: 4, SEQ ID NO: 7, and SEQ ID NO: 8 comprising:
- (i) contacting an abnormal cell sample from a subject with a cytolytic T cell specific for said complexes, and
- (ii) determining lysis of at least part of said abnormal cell sample as an indication of a candidate for said treatment.
- 2. The method of claim 1, wherein said MHC molecule is HLA-A2.
- 3. The method of claim 1, wherein said MHC molecule is HLA-B44.
- 4. Method for treating a subject with a cellular abnormality, comprising administering to said subject an amount of an agent which provokes a cytolytic T cell response to cells presenting complexes of an MHC molecule and a tyrosinase derived peptide selected from the group consisting of SEQ ID NO: 4, SEQ ID NO: 7, and SEQ ID NO: 8 on their surfaces sufficient to provoke a response to abnormal cells presenting said complexes on their surfaces.
- 5. The method of claim 3, wherein said MHC molecule is HLA-A2.
- 6. The method of claim 4, wherein said MHC molecule is HLA-B44.
- 7. The method of claim 4, wherein said agent comprises a vector which codes for human tyrosinase.
- 8. The method of claim 7, wherein said agent further comprises a second vector which codes for HLA-A2.
- 9. The method of claim 7, wherein said agent further comprises a second vector which codes for HLA-B44.
- 10. The method of claim 7, wherein said vector also codes for HLA-A2.
- 11. The method of claim 7, wherein said vector also codes for HLA-B44.

- 12. The method of claim 4, wherein said agent is a sample of non-proliferative cells which present said complexes on their surfaces.
- 13. Method for treating a cellular abnormality comprising administering to a subject with a cellular abnormality characterized by presentation of complexes of an MHC molecule and a tyrosinase derived peptide selected from the group consisting of SEQ ID NO: 4, SEQ ID NO: 7 and SEQ ID NO: 8 on surfaces of abnormal cells an amount of cytolytic T cells specific for said complexes sufficient to lyse said abnormal cells.
- 14. The method of claim 13, wherein said MHC molecule is HLA-A2.
- 15. The method of claim 13, wherein said MHC molecule is HLA-B44.
- 16. The method of claim 13, wherein said cytolytic T cells are autologous.
- 17. Isolated cytolytic T cell specific for a complex of an MHC molecule selected from the group consisting of HLA-A2 and a tyrosinase derived peptide selected from the group consisting of SEQ ID NO: 4, SEQ ID NO: 7, and SEQ ID NO: 8.
- 18. The isolated cytolytic T cell of claim 17, specific for a complex of HLA-A2 and SEQ ID NO: 4.
- 19. The isolated cytolytic T cell of claim 17, specific for a complex of HLA-B44 and SEQ ID NO: 7.
- 20. The isolated cytolytic T cell of claim 17, specific for a complex of HLA-B44 and SEQ ID NO: 8.
- 21. Method for identifying an abnormal cell which presents a complex of an MHC molecule and a tyrosinase derived peptide selected from the group consisting of SEQ ID NO: 4, SEQ ID NO: 7, and SEQ ID NO: 8 on its surface comprising contacting a sample of abnormal cells with a cytolytic T cell specific for said complex and determining lysis of said abnormal cells as a determination of cells which present said complex.
- 22. The method of claim 21, wherein said MHC molecule is HLA-A2.

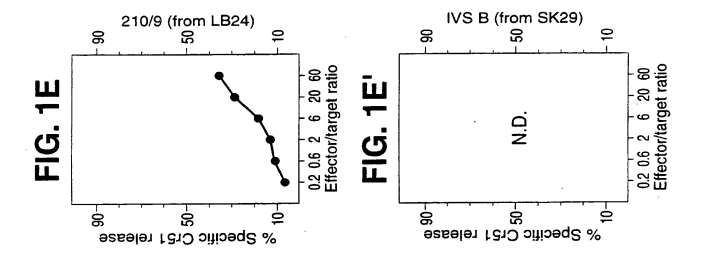
- 23. The method of claim 21, wherein said MHC molecule is HLA-B44.
- 24. Isolated peptide selected from the group consisting of SEQ ID NO: 4, SEQ ID NO: 7, and SEQ ID NO: 8.

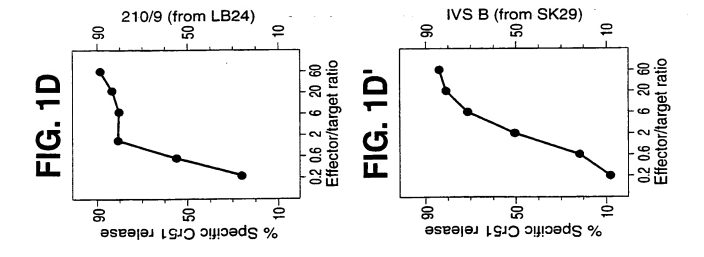
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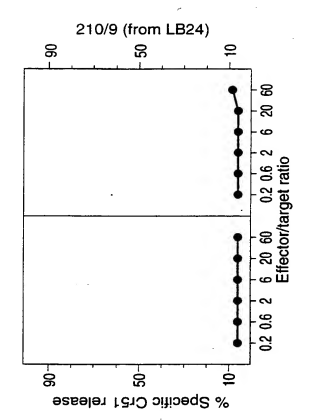




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FIG. 1F



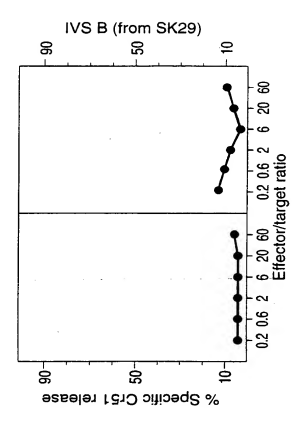
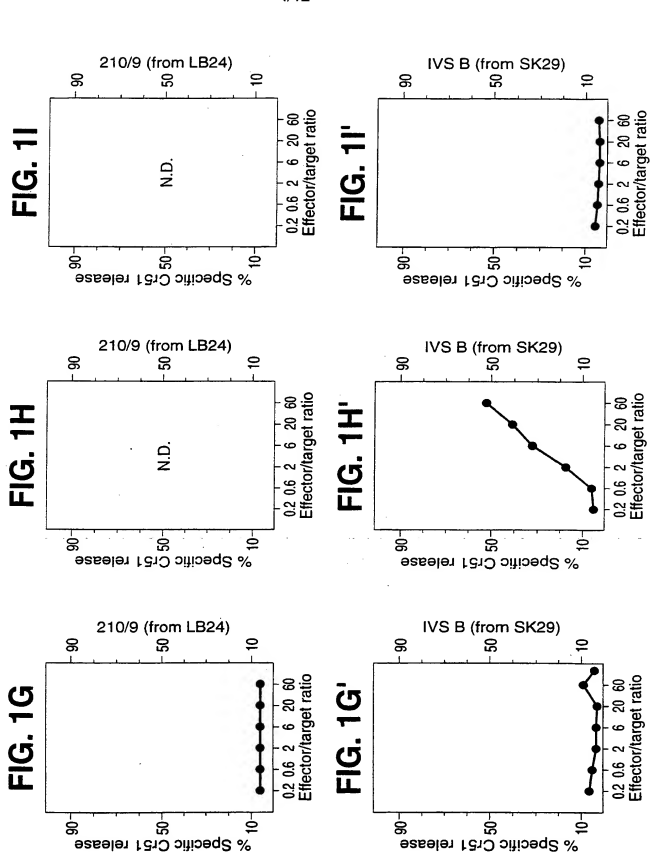


FIG. 1F

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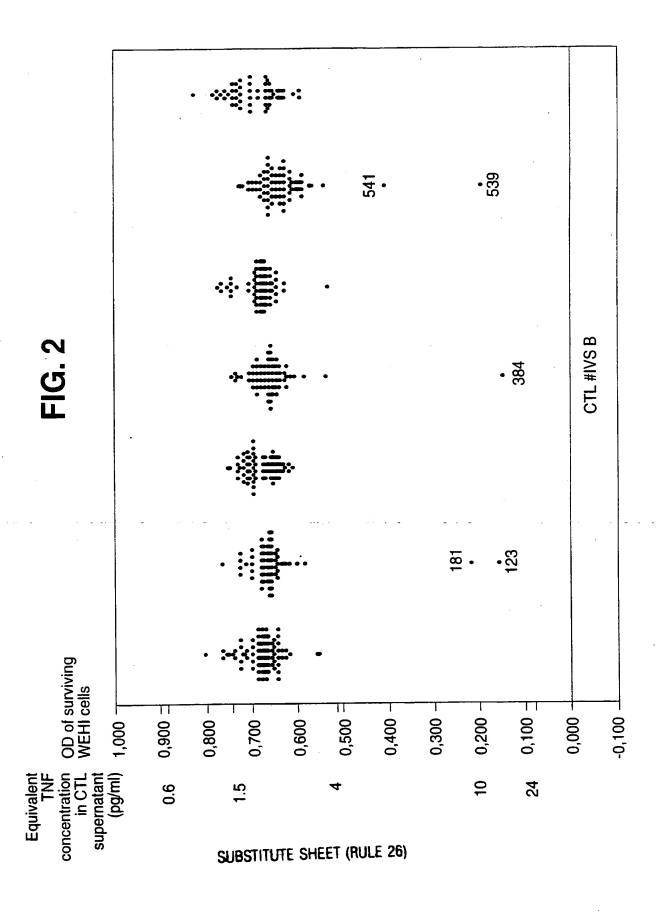


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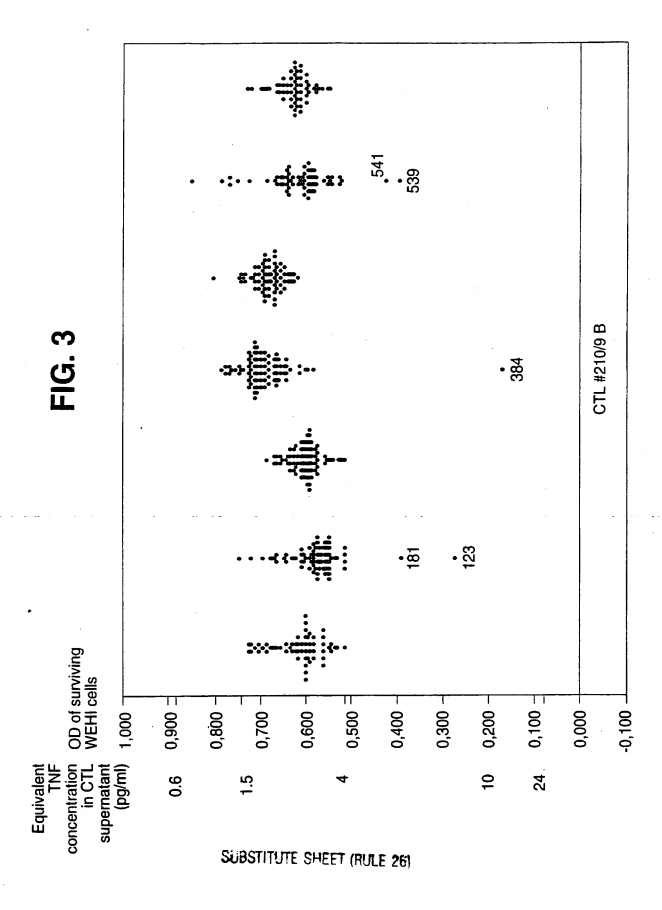
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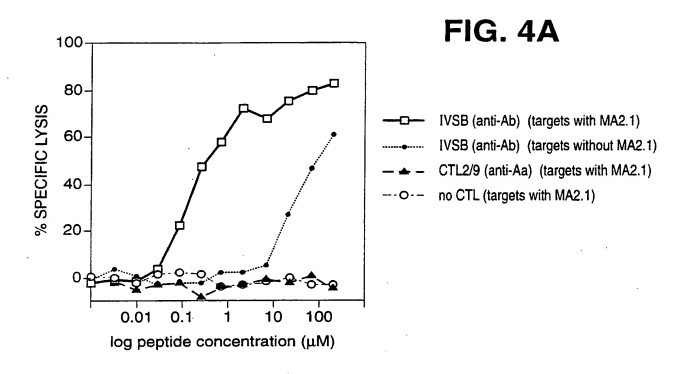


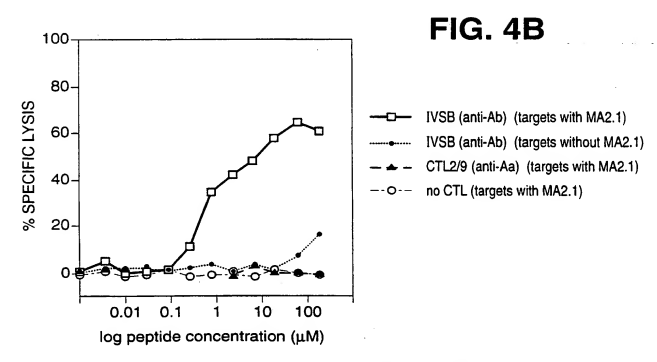
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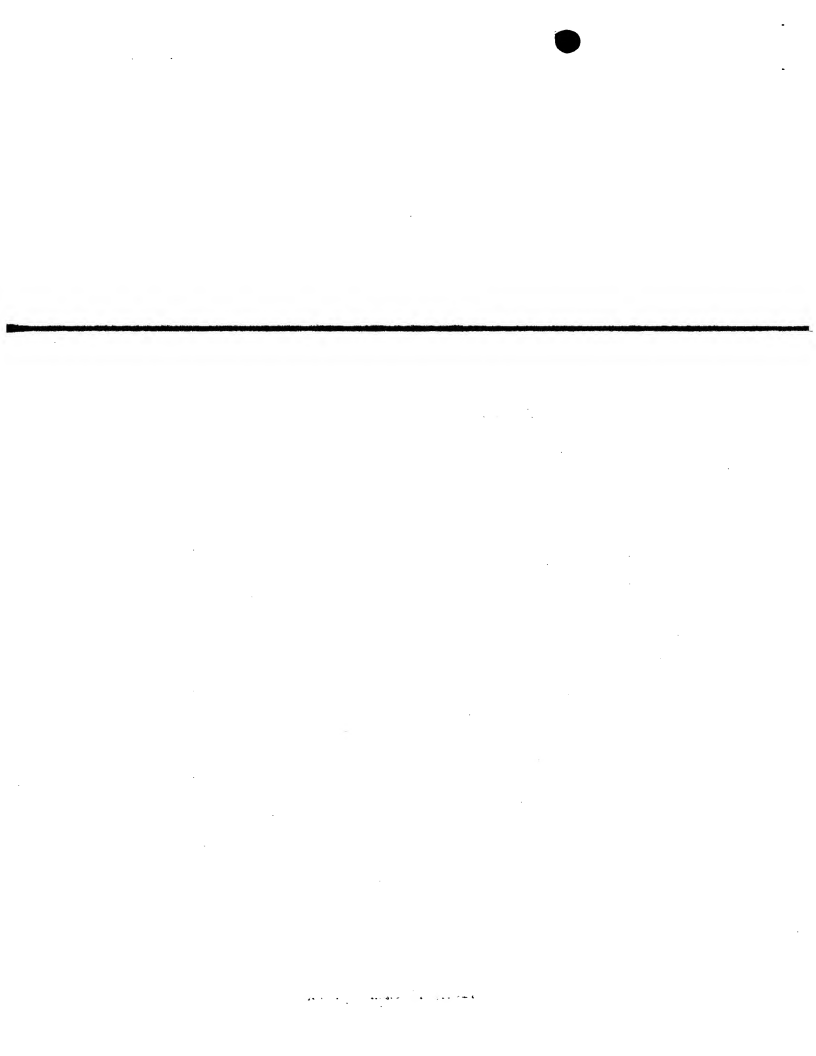
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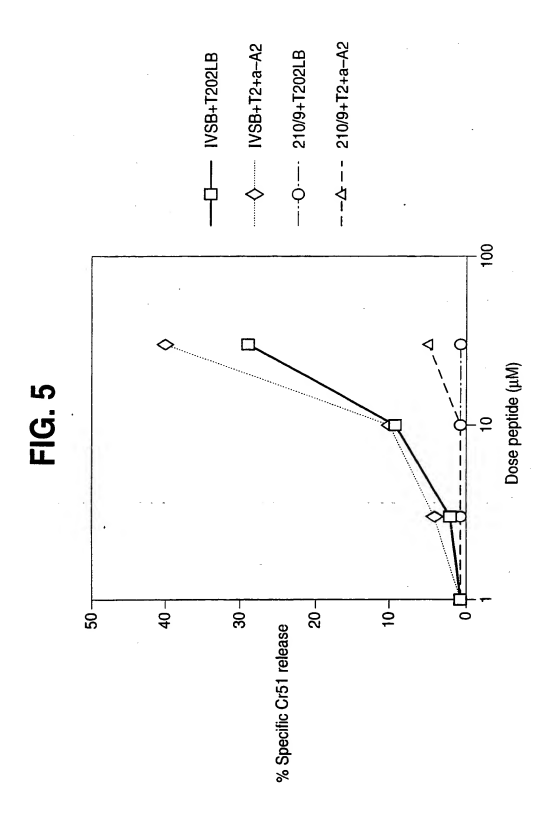
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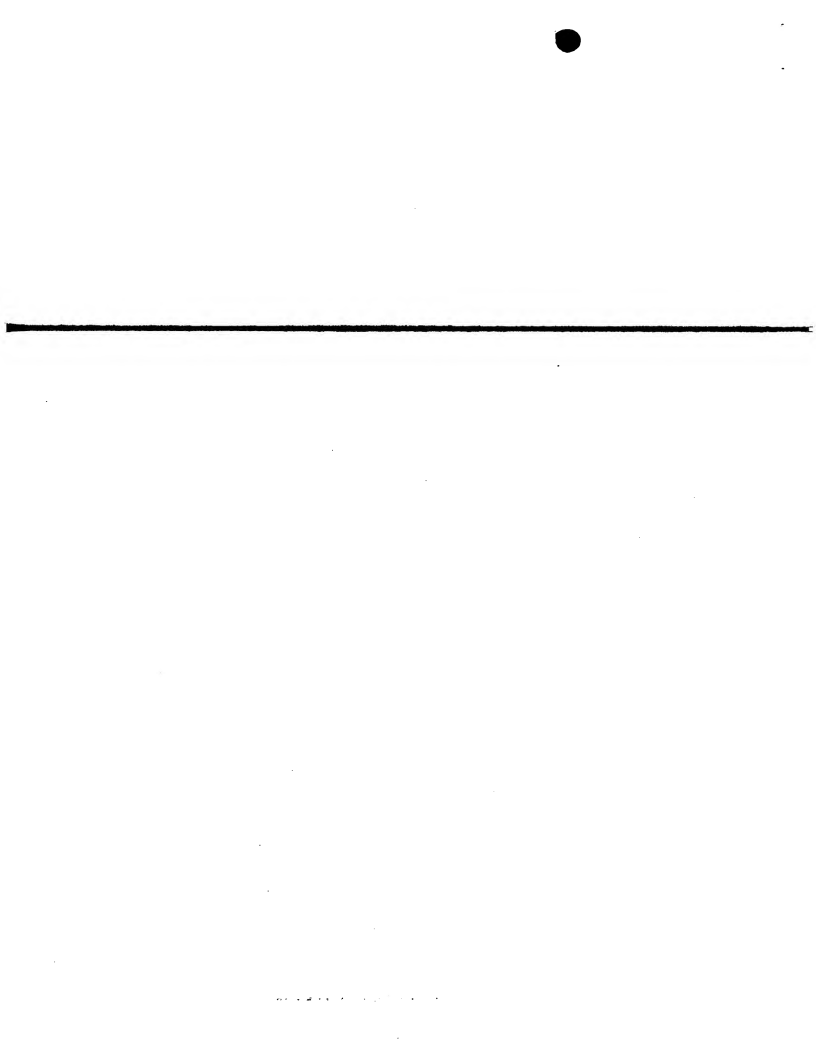


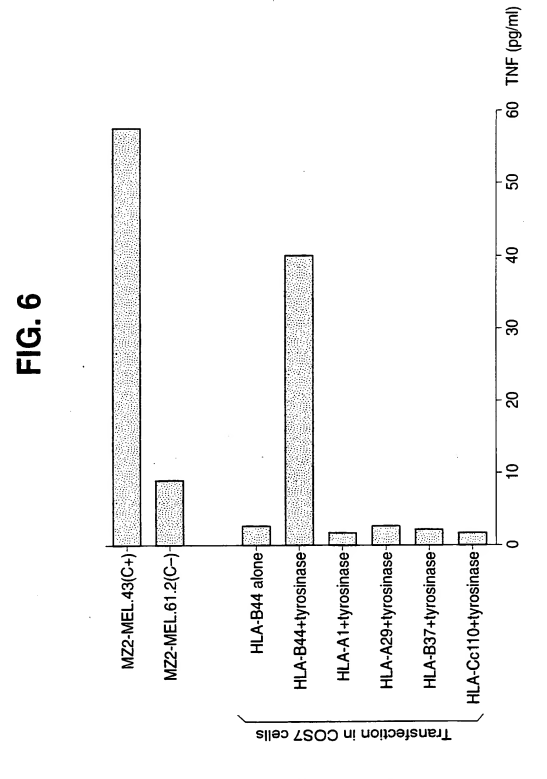
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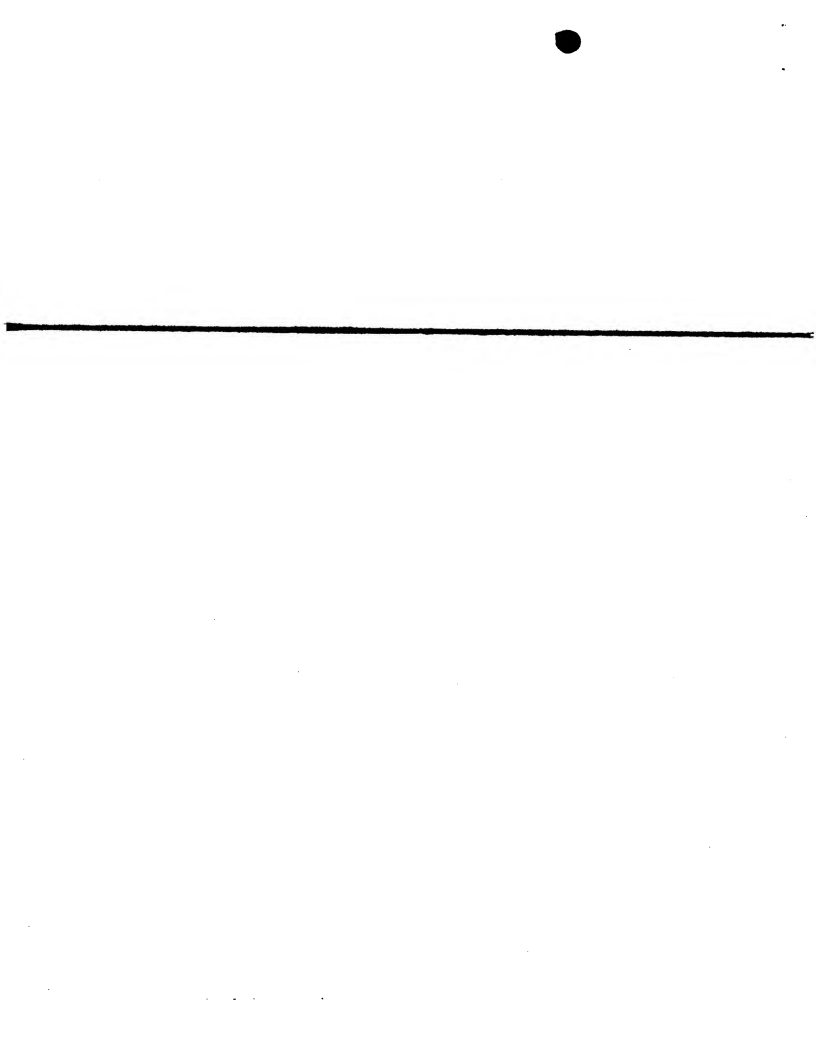


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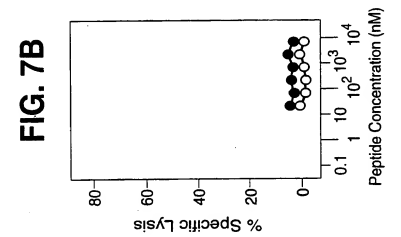
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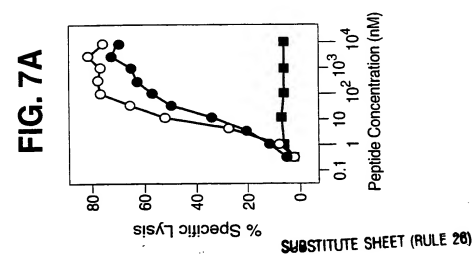
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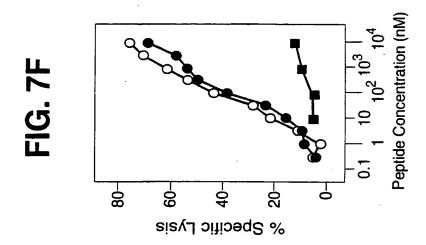
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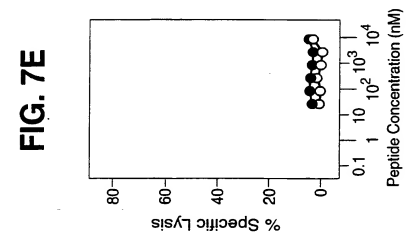


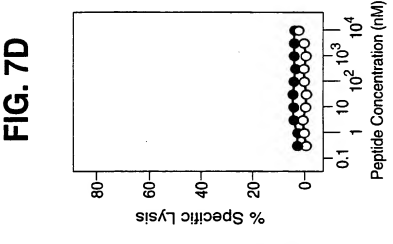


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FIG. 8A

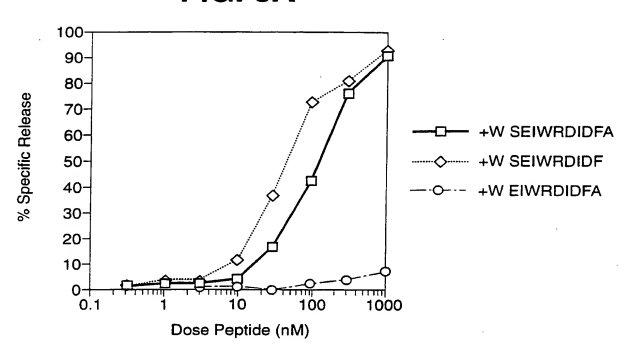
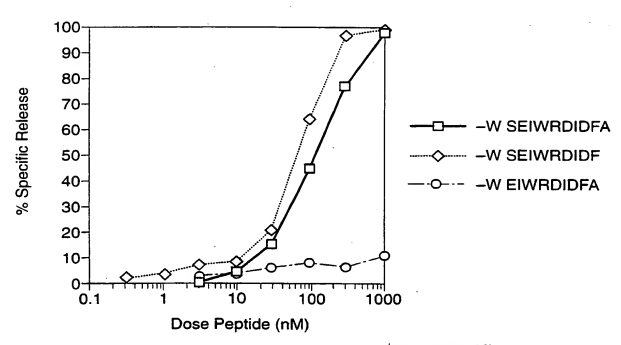


FIG. 8B



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International application No.
PCT/US95/01990

IPC(6) US CL	SSIFICATION OF SUBJECT MATTER :C12Q 1/04; C12N 5/08; C07K 7/06 :435/7.24, 240.2; 530/328 to International Patent Classification (IPC) or to both a	national classification and IPC				
	LDS SEARCHED					
Minimum d	locumentation searched (classification system followed	by classification symbols)				
	424/ 93.21; 435/7.23, 7.24, 240.2; 514/ 15, 16; 530					
Documenta	tion searched other than minimum documentation to the	extent that such documents are included	in the fields searched			
Electronic o	data base consulted during the international search (na	me of data base and, where practicable,	search terms used)			
1	ALOG, SEQUENCE SEARCH (SEQ ID NOs: 4, 7,					
C. DOC	CUMENTS CONSIDERED TO BE RELEVANT	·				
Category*	Citation of document, with indication, where ap	propriate, of the relevant passages	Relevant to claim No.			
X,P	EUROPEAN JOURNAL OF IMMUN	IOLOGY, Vol. 24, issued	24			
	March 1994, Wolfel et al., "Two					
Y,P	recognized on HLA-A2 melanomas lymphocytes," pages 759-764, see		1-3, 17-23			
Y	I. ROITT et al., "Immunology, 3rd	Edition." published 1993	1-3, 17-18, 21-			
•	by Mosby (St. Louis, Mo, USA)		24			
	paragraph bridging pages 6.10-6.1					
Υ	JOURNAL OF EXPERIMENTAL ME	DICINE, Vol. 178, issued	1-3, 17-24			
	August 1993, Brichard et al., "The		•			
	for an Antigen Recognized by					
	Lymphocytes on HLA-A2 Melanom	nas," pages 489-495, see				
	entire document.					
}						
		•				
X Furt	her documents are listed in the continuation of Box C					
1	pecial categories of cited documents:  comment defining the general state of the art which is not considered	"T" later document published after the inte date and not in conflict with the applic	ation but cited to understand the			
to	be of particular relevance	principle or theory underlying the inv  "X" document of particular relevance; th	_			
	rifer document published on or after the international filing date	considered novel or cannot be considered when the document is taken alone				
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	actual completion of the international search	Date of mailing of the international see	arch report			
08 JUNE	. 1995					
Name and	mailing address f the ISA/US oner of Patents and Trademarks	Authorized officer  ROBERT D. BUDENS	Farnos			
Box PCT	on, D.C. 20231	ROBERT D. BUDENS	- 1200-4			
	No. (703) 305-3230	Telephone No. (703) 308-0196				

International application No. PCT/US95/01990

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim N
Y	INTERNATIONAL JOURNAL OF CANCER, Vol. 55, issued 1993, Wolfel et al., "Analysis of Antigens Recognized On Human Melanoma Cells By A2-Restricted Cytolytic T Lymphocytes (CTL)," pages 237-244, see entire document.	1-3, 17-24
?	JOURNAL OF IMMUNOTHERAPY, Vol. 14, issued 1993, Coulie et al., "Genes Coding For Tumor Antigens Recognized By Human Cytolytic T Lymphocytes," pages 104-109, see entire document.	1-3, 17-24
<b>′</b> ,P	CANCER RESEARCH, Vol. 54, issued 15 June 1994, Robbins et al., "Recognition of Tyrosinase By Tumor-Infiltrating Lymphocytes from a Patient Responding to Immunotherapy, "pages 3124-3126, see entire document.	1-3, 17-24
	THE JOURNAL OF IMMUNOLOGY, Vol. 150, No. 7, issued 01 April 1993, Slingluff, Jr. et al., "Recognition of Human Melanoma Cells by HLA-A2.1-Restricted Cytotoxic T Lymphocytes Is Mediated by at Least Six Shared Peptide Epitopes," pages 2955-2963.	1-3, 17-24
	THE JOURNAL OF IMMUNOLOGY, Vol. 154, issued 1995, Visseren et al., "CTL Specific for the Tyrosinase Autoantigen Can Be Induced from Healthy Donor Blood to Lyse Melanoma Cells," pages 3991-3998, see entire document.	1-3, 17-24
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International application No. PCT/US95/01990

Box 1 Observations where certain claims were f und unsearchable (Continuati n of item 1 f first sheet)
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. Claims Nos.:  because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
Please See Extra Sheet.
1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. X As only some of the required additional search fees were timely paid by the applicant, this international search report cov rs only those claims for which fees were paid, specifically claims Nos.:  1-3, 17-24
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark n Protest
N protest accompanied the payment f additi nal search fees.

International application No. PCT/US95/01990

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be examined, the appropriate additional examination fees must be paid.

Group I, claims 1-2, 17-18, 21-22 and 24, drawn to methods of identifying candidates for therapy.

Group II, claims 4-12, drawn to a second method, methods of treatment using a therapeutic agent.

Group III, claims 13-16, drawn to a third method, methods of treatment using cytotoxic T lymphocytes.

Group IV, claim 24, drawn to isolated tyrosinase peptides.

The inventions listed as Groups I-III and IV do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they tack the same or corresponding special technical features for the following reasons: The inventions of Groups I-III are directed to distinct methods using different method steps and uses. Further, the product of Group IV is independent of the methods encompassed in the inventions of Groups I-III. The product of Group IV does not share a special technical feature with the methods of Groups I-III.

This application contains claims directed to more than one species of the generic invention. These species are deemed to lack Unity of Invention because they are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for more than one species to be examined, the appropriate additional examination fees must be paid. The species are as follows:

Group V, claims 3, 19-20 and 23, drawn to a second HLA haplotype, HLA-B44.

Group VI, claims 1-3 and 17-24, drawn to a second species of tyrosinase peptide, SEQ ID NO: 7.

Group VII, claims 1-3 and 17-24, drawn to a third species of tyrosinase peptide, SEQ ID NO: 8.

The following claims are generic: 1, 21

The species listed above do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, the species lack the same or corresponding special technical features for the following reasons: Group V is directed to a different HLA haplotype distinct from HLA-A2 and differing in structure and function from HLA-A2. Further, the species of Groups VI-VII are directed to tyrosinase peptides differing in their primary amino acid sequence, structure and physical properties and are distinct from the tyrosinase peptide of SEQ ID NO:4. Groups V-VII do not share a special technical feature.

Accordingly, the claims are not so linked by a special technical feature within the meaning of PCT Rule 13.2 so as to form a single inventive concept.